Developmental Sculpting of Intracortical Circuits by MHC Class I H2-Db and H2-Kb

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Abstract

Synapse pruning is an activity-regulated process needed for proper circuit sculpting in the developing brain. Major histocompatibility class I (MHC1) molecules are regulated by activity, but little is known about their role in the development of connectivity in cortex. Here we show that protein for 2 MHC1 molecules H2-Kb and H2-Db is associated with synapses in the visual cortex. Pyramidal neurons in mice lacking H2-Kb and H2-Db (KbDb KO) have more extensive cortical connectivity than normal. Modified rabies virus tracing was used to monitor the extent of pyramidal cell connectivity: Horizontal connectivity is greater in the visual cortex of KbDb KO mice. Basal dendrites of L2/3 pyramids, where many horizontal connections terminate, are more highly branched and have elevated spine density in the KO. Furthermore, the density of axonal boutons is elevated within L2/3 of mutant mice. These increases are accompanied by elevated miniature excitatory postsynaptic current frequency, consistent with an increase in functional synapses. This functional and anatomical increase in intracortical connectivity is also associated with enhanced ocular dominance plasticity that persists into adulthood. Thus, these MHC1 proteins regulate sculpting of local cortical circuits and in their absence, the excess connectivity can function as a substrate for cortical plasticity throughout life.

Key words: activity-dependent development, horizontal connections, major histocompatibility complex class I, modified rabies circuit tracing, visual cortex

Introduction

Intracortical circuits are sculpted by neural activity and experience during developmental critical periods (Katz and Shatz 1996), and retain a more limited ability to remodel during maturity (Knudsen 2004). In many binocular animals, L2/3 and L5 pyramidal neurons in visual cortex extend long horizontal axons laterally up to a few millimeters away as well as across the corpus callosum, connecting functionally relevant subsets of neurons to each other (Livingstone and Hubel 1984; Gilbert and Wiesel 1989; Gilbert 1992; Kenan-Vaknin et al. 1992; Salin and Bullier 1995; Trachtenberg and Stryker 2001; Tanigawa et al. 2005; Van Hooser et al. 2006). Initially, in the visual cortex, these axons are exuberant (Callaway and Katz 1990). During the critical period, these horizontal connections are pruned in a process requiring visual experience (Callaway and Katz 1991) and neural activity (Gilbert and Wiesel 1989; Callaway and Katz 1990; Lund et al. 1993; Malach et al. 1993; Durack and Katz 1996; Ruthazer and Stryker 1996). Dendrites and spines in many regions of the brain also undergo an activity-regulated period of growth, followed by retraction (Cline 2001; Sin et al. 2002; Vaillant et al. 2002; Hofer et al. 2006a, 2006b; De Marco Garcia et al. 2011; Chen et al. 2012). In the adult, anatomical changes in horizontal connections in...
dependent synaptic plasticity. When one eye is closed or removed, the deprived eye loses the ability to activate cortical neurons, which become more responsive to the open eye (Wiesel and Hubel 1963; Hubel et al. 1977). During the critical period, visual deprivation causes a strong shift in the OD of cortical neurons toward the open eye (Hubel et al. 1977; Shatz 1990; Gordon and Stryker 1996; Frenkel and Bear 2004). In adult visual cortex, OD plasticity can be elicited but requires significantly longer periods of monocular deprivation (MD; Tagawa et al. 2005; Sato and Stryker 2008). Many studies have identified molecular mechanisms that contribute to activity-dependent synaptic plasticity in the visual cortex (Hensch 2005; Flavell and Greenberg 2008; Tropea et al. 2009; Levelt and Hübener 2012), but the link from candidate molecules to experience-dependent structural change has not been explored as extensively.

One set of candidate molecules that could link activity-dependent structural changes to alterations in synaptic and circuit-level plasticity are the major histocompatibility class I (MHC-I) genes, which were discovered in a screen for activity-regulated genes in the developing visual system (Corriu et al. 1998; Shatz 2009; Elmer and McAllister 2012). In the developing lateral geniculate nucleus (LGN) of the thalamus, neural activity is known to be required for synapse pruning and eye-specific layer formation (Huberman et al. 2008). Just 2 of the over 50 MHC-I genes, H2-Kb and H2-Db, are known to regulate synapse pruning via alterations in synaptic learning rules and glutamate receptor subunit composition at retinogeniculate synapses (Lee et al. 2014). Mice lacking both H2-Kb and H2-Db (KbDb KO) have less synapse elimination compared with wild type (WT), and elimination can be restored to WT levels by selectively expressing H2-Db in LGN neurons (Lee et al. 2014). H2-Kb and H2-Db are also both expressed in cortical neurons including pyramidal cells (Huh et al. 2000; Datwani et al. 2009). During the critical period in the visual cortex, KbDb KO mice have enhanced OD plasticity, as assessed by measuring open eye strengthening following MD (Datwani et al. 2009; Shatz 2009). In vitro studies on β2m/Tap1 KO mice, which lack surface expression of the vast majority of MHC-I proteins, reveal an increase in synapse density, while over expression of H2-Kb in vitro resulted in a decrease in synapse density (Goddard et al. 2007; Glynn et al. 2011); both observations are consistent with the idea that these MHC-I proteins contribute to synapse pruning. Here, we examined whether the specific MHC-I molecules H2-Kb and H2-Db also regulate developmental sculpting of intracortical connectivity in vivo in the visual cortex.

Materials and Methods

Animals

KbDb KO mice, offspring of breeding pairs on a C57BL/6 background, were originally generously provided by H. Ploegh and are now maintained in our colony (Cambridge, MA, USA; Vugmeyster et al. 1998). C57BL/6 (i.e., KbDb WT) controls were purchased (Charles River) or bred in Stanford facilities. For modified rabies virus circuit tracing experiments, CaMK2α-CreER; CAG-stop-tTA2 double transgenic mice were provided by L. Luo (Miyamichi et al. 2011; Stanford, CA, USA) and bred to KbDb KO mice. F1 generation CaMK2α-CreER; CAG-stop-tTA2; KbDb+/− mice were bred to KbDb−/− mice to produce F2 experimental animals (CaMK2α-CreER; CAG-stop-tTA2; KbDb−/− or KbDb−/−), control animals (CAG-stop-tTA2; KbDb−/− or KbDb−/−), and breeders (CaMK2α-CreER; CAG-stop-tTA2; KbDb−/−). All mice were maintained in a pathogen-free environment. All experiments using animals were performed blind to the genotype and in accordance with a protocol approved by the Stanford University animal care and use committee and in keeping with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

Retrograde Circuit Tracing with Modified Rabies Virus

See Figure 1A and Supplementary Figure 1 for diagrams and complete description of methods, respectively. All handling of rabies virus followed procedures approved by Stanford University’s Administrative Panel on Biosafety (APB) for Biosafety Level 2. pAAV2 TRE-histone-mCherry-2a-TVA-2a-RG (TRE-mCTG) was constructed by replacing GFP-coding sequence of TRE-HTG (Miyamichi et al. 2011) with that of mCherry by using PCR-based cloning. Recombinant adeno-associated virus (AAV; serotype 2) was produced in the University of North Carolina viral core. Mice were injected IP with 0.1 mg Tamoxifen Free Base (Sigma, T5648) suspended in 2% ethanol and 98% corn oil (Sigma, 63156) at P10–P14. At P21–P23, mice were injected with 150 nL of AAV2 TRE-mCTG. Mice were placed in a Kopf stereotaxic set-up under approximately 1–2% isoflurane inhalable anesthesia. AAV was injected using a NanoFil 34–36 gauge beveled syringe (NanoFil-100, World Precision Instruments) at 50 nL/min into visual cortex through a small drill hole (coordinates: +2.25 mm lateral, 3.4 mm posterior to Bregma, −0.5 mm depth from brain surface). Fourteen days after AAV injection (P35–P37), mice were injected at 50 nL/min with 150 nL rabies-DG-GFP + EnvA [prepared according to the protocol (Osakada and Callaway 2013) or purchased from Salk Institute Viral Core], through the same drill hole as for AAV injection. Note that the volumes of injected virus were small (150 nL) to restrict the number of starter cells and the location of starter cells to a local region of visual cortex, permitting connections within visual cortex to be analyzed. In addition, to validate this retrograde tracing method, which previously had only been used for long distance connectivity (Miyamichi et al. 2011), in one additional experiment, much larger injections were made (1000 nL of each virus) and retrograde labeling in LGN and contralateral visual cortex was assessed. Five days after the modified rabies injection (P40–P42), animals were deeply anesthetized and perfused with ice-cold PBS and 4% paraformaldehyde. Coronal sections of visual cortex were cut at 50 μm thickness using a Leica VT1200s vibratome (Leica Microsystems, Inc., IL, USA) and mounted on slides with ProLong Gold Anti-Fade Reagent (Life Technologies, CA, USA).

Horizontal Connectivity Analysis

The number of “starter cells” was counted for each analyzed animal, where a starter cell is defined as a neuron infected by both AAV and modified rabies virus, determined by co-localization of mCherry and GFP (Fig. 1A and Supplementary Fig. 1). The number of GFP-labeled long-range presynaptic partners was counted, defined as cells falling outside of 250 μm laterally from the injection site. Total connectivity was determined by counting all GFP-labeled cells, dividing by the number of starter cells, and subtracting the average background number of GFP-labeled cells observed in control Cre− mice. Counting of GFP-labeled cells was initiated in the furthest posterior coronal section of visual cortex containing a starter cell and all subsequent anterior sections of cortex.
Dendritic Complexity and Spine Density Analyses

Dendritic complexity was assessed using Sholl analysis (Sholl 1953). Isolated GFP-labeled excitatory pyramids in L2/3 were imaged using a Prairie 2-photon microscope for dendritic morphology analysis. Cells were identified as excitatory based on morphology, and imaged if at least 75 μm continuous basal and/or apical dendrites proximal to the cell body were in the plane of sectioning. We focused specifically on an analysis of proximal dendrites because these receive the bulk of intracortical horizontal connections from other pyramidal cells (Gilbert and Wiesel 1989; Thomson and Bannister 2003; Feldmeyer et al. 2006). Imaging was carried out with a ×63 water immersion objective corrected for slide thickness, and a ×1 zoom for full cell imaging, and a ×4 zoom for dendrite imaging (51 μm × 51 μm field of view). Dendritic morphology was analyzed by reconstructing cells with Fiji’s “Simple Neurite Tracer” followed by Fiji’s Sholl analysis of dendrites belonging to identified pyramidal neurons in 50-μm thick visual cortex slices.

Axon Bouton Analysis

Biotinylated dextran amine (BDA) was used for anterograde labeling of horizontal axonal projections in L2/3 of visual cortex (as described by Veenman et al. 1992). Stereotaxic surgeries were performed on a Kopf stereotaxic set-up. At P26 or P27, mice were anesthetized with approximately 2.5% isoflurane and a small hole was drilled into the skull in visual cortex (relative to Bregma: X: −3.4 mm, Y: −2.25 mm, Z: −0.5 mm). A small needle was used to pierce the dura, and 200 nL of BDA (10 000 MW, NeuroTrace BDA-10 000 Neuronal Tracer Kit, Molecular Probes, Eugene, OR, USA) was injected into each site using a NanoFil syringe (World Precision Instruments, Inc., Sarasota, CA, USA). After 14 days (P40–P41), mice were deeply anesthetized and perfused, brains were removed and 50 μm coronal sections were cut using a vibratome (VT 1000 S; Leica Microsystems, Wetzlar, Germany). Sections were incubated with 0.6 μg/mL of avidin–horse-radish peroxidase (NeuroTrace BDA-10 000 Neuronal Tracer Kit) and reacted with 1% diaminobenzidine to visualize tracer, and mounted on slides. Sections were imaged at ×10 using bright-field microscopy (Nikon Eclipse E800, Melville, NJ, USA). The density of en passant boutons in visual cortex ipsilateral to the injection site was assessed using ImageJ.

Visual Cortex Slice Physiology

Cortical slices and physiology were carried out as in Goddard et al. (2007) and Lee et al. (2014). After brief intracardinal perfusion of ice-cold ACSF (in mM; 125 NaCl, 26 NaHCO3, 2.3 KCl, 1.26 KH2PO4, 1.3 MgCl2, 2.5 CaCl2, and 10 glucose, aerated with 95% O2/5% CO2), brains from P26 to P31 male mice were removed and coronal sections (400 μm) including visual cortex were made using a vibratome (Leica VT1000S, Leica Microsystems, Inc., IL, USA) in ice-cold NMDG solution (in mM: 135 NMDG, 1 KCl, 1.2 KH2PO4, 1.5 MgCl2, 0.5 CaCl2, 20 choline bicarbonate, 1.25 Glu, 10 D-glucose, 10 NaHCO3, 1 CaCl2, 0.5 KCl, 0.1 Na2ATP, 1.25 NaH2PO4, 0.4 MgSO4). Sections were stored in ice-cold ACSF and transferred to a recording chamber. Target areas were identified with an infrared differential interference contrast (DIC) microscope (Leica DMI6000).
and 10 glucose). Sections were transferred to a recovery chamber containing ACSF at 37 °C for 30 min and at room temperature for an additional 30 min before recordings. Whole-cell patch-clamp recordings were performed from pyramidal neurons in layer 2/3 of visual cortex. Recording pipette (2–4 MΩ) contained Cs+-based internal solution (in mM: 105 CsCl, 20 TEA-Cl, 2 MgCl₂, 1 EGTA, 10 HEPES, 3 Mg₂-ATP, 15 phosphocreatine, 1 Na-GTP, 5 QX-314, pH 7.4, and 280 mOsm). Miniature excitatory postsynaptic currents (mEPSCs) were isolated by applying TTX (1 µM, Sigma), SR95531 (20 µM; Tocris) to block GABA_A receptors, and APV (100 µM, Tocris) to block NMDA receptors. All recordings were done at 30–32 °C in a chamber with constant ACSF flow. Synaptic responses were recorded using an Axopatch 200B amplifier (Molecular Devices, CA, USA), digitized using Digidata 1222A (Axon Instruments, CA, USA), and data acquisition was performed by Clampex 9.2 (Axon Instrument, CA, USA). Data analysis was conducted using the MiniAnalysis software (ver. 6.0.7, Synaptosoft).

Synaptosome Preparation and Western Blots

Synaptosome preparations and western blots for H2-Kb and pan-KbDb/KbDb were carried out as detailed in Adelson et al. (2012). Synaptosome-enriched fractions were prepared as described (Johnson et al. 1997; Yin et al. 2002). Briefly, freshly isolated brain hemispheres were homogenized by 12 strokes of a Dounce homogenizer in homogenization buffer (10 mM HEPES pH 7.3, 0.5 mM EGTA, 33% sucrose, 4 mM Pefabloc SC PLUS (Roche), and 0.2 mM phenylmethylsulfonyl fluoride), and centrifuged for 10 min at 2000 × g. Supernatants were passed through three-layered 100 µm pore nylon membranes, and then passed through 5 µm nitrocellulose filters. For further purification to synaptosomes, filtrates were centrifuged at 10,000 × g for 10 min, and pellets were resuspended in homogenization buffer and SDS-PAGE sample buffer. Antibodies against H2-Kb and H2-Kb/KbDb were used at 1:500 dilution (Adelson et al. 2012; Lee et al. 2014). Samples were then electrophoresed on an SDS-PAGE gel, transferred to the Immobilon-P PVDF transfer membrane (Millipore), and western blotted with rabbit monoclonal antibodies to H2-Kb or H2-D/ Kb, at a dilution of 1:500.

Enucleations and Arc mRNA Induction

Enucleations and Arc mRNA induction were performed as detailed in Tagawa et al. (2005). Briefly, KbDb KO and WT controls were monocularly enucleated at P100 (monocular enucleation treatment group) or P109 (normally reared treatment group) using 2–3% isoflurane anesthesia. Eyelids were sealed with Vetbond tissue adhesive glue (3M, St Paul, MN, USA). All mice were placed in total darkness for 16 h overnight at P109. At P110, mice were exposed to bright light for 30 min, followed by anesthesia with 3% isoflurane and immediate brain removal. Brains were embedded in Thermo Scientific’s M-1 Embedding Matrix for frozen sectioning and frozen using 100% ethanol and dry ice, stored at 80 °C, and sectioned at 16 µm on a Thermo Scientific Shandon cryostat.

Isotopic In Situ, Autoradiography, and Analysis

Isotopic in situ hybridization for Arc mRNA was performed as previously described (Lein and Shatz 2000; Tagawa et al. 2005) using an isotopic (5S²) probe that hybridized to Arc mRNA (generated from full-length mouse Arc cDNA, gift of P. Worley, Johns Hopkins University, Lyford et al. 1995). Autoradiography in photographic emulsion was allowed to develop for 2–3 days before development. Analysis of signal was accomplished using the NeuroLens software (version 1.7.3, Massachusetts General Hospital, Université de Montréal et CRIUGM; neurolems.org).

Results

Modified Rabies Virus Circuit Tracing Reveals Greater Density of Intracortical Pyramidal Neuron Connectivity in KDbb KO Visual Cortex

To address whether H2-Kb and H2-Db regulate patterns of horizontal connectivity in cortex, we employed a modified rabies virus tracing method. By using a combined transgenic and viral vector approach, modified rabies virus circuit tracing can identify a small subset of starter cells, as well as their presynaptic partners that get labeled transsynaptically via rabies virus infection, which crosses the synapse retrogradely from the starter cells (Fig. 1A; Supplementary Fig. 1; Wickersham et al. 2007; Wall et al. 2010; Miyamichi et al. 2011). This method was initially developed to examine long-range patterns of connectivity in the olfactory system (Miyamichi et al. 2011). Here, we employ it in a new application to examine intracortical patterns of connectivity in WT and KbDb KO mice within a single cerebral cortical area.

To study connectivity of the CaMK2α* pyramidal neurons primarily responsible for the horizontal connections in visual cortex (Gilbert and Wiesel 1989; Thomson and Bannister 2003; Feldmeyer et al. 2006), WT or KbDb KO mice were generated on a transgenic background needed for rabies tracing. A transgenic line (Tg) carrying ubiquitous expression of a tetracycline transactivator element (tTA2) under transcriptional control of a fixod stop cassette (CAG-stop-tTA2), as well as Cre fused to an estrogen receptor with a CaMK2α promoter (CaMK2α-CreER; Miyamichi et al. 2011), was used (see Materials and Methods); this line, henceforward called the transgenic line (Tg), was crossed to WT and KbDb KO mice, generating WT-Tg and KbDb KO-Tg mice used here. In these mice, tamoxifen administration (P10–P14) activates Cre, inducing the expression of tTA2 only in CaMK2α* pyramidal neurons. One week later, when tTA2 is well expressed (Miyamichi et al. 2011), an AAV was stereotaxically injected into visual cortex under the control of a tetracycline-response element (TRE), causing the production of histone-mCherry as well as the avian receptor TVA (Barnard et al. 2006), and rabies glycoprotein (RG) in infected CaMK2α* cells. Two weeks later at P35, a modified rabies virus expressing GFP and the avian envelope protein EnvA (which recognizes and selectively infects TVA* cells) was stereotaxically injected into the same region of visual cortex. Cells infected with both AAV and modified rabies virus were termed “starter cells,” identified by co-labeling of histone-mCherry and GFP. The AAV-provided RG protein (needed for synapse jumping) allowed the rabies virus to travel retrogradely to presynaptic partner cells (GFP-labeled only). Cortical connectivity in WT-Tg and KbDb KO-Tg was assessed 5 days after rabies infection, at P40.

To validate methodology in WT-Tg mice, we first demonstrated the presence of AAV-infected histone-mCherry-labeled cells, modified rabies-infected GFP-labeled cells, and the “starter cells” in which both fluorophores are colocalized, rendering them yellow (Fig. 1B). A low dose of tamoxifen that did not cause significant body weight change (Materials and Methods; Supplementary Fig. 2A) was used to activate Cre. At the site of the injection, control mice lacking Cre never had AAV-infected mCherry-labeled cells, and only a low level of rabies-infected GFP-labeled cells, likely due to slight leakiness of the TRE promoter (Miyamichi et al. 2011). The number of GFP-labeled cells
KbDb KO L2/3 Pyramidal Cells Have More Complex Basal Dendrites and Elevated Spine Density

Pyramidal neurons in L2/3 and L5 of visual cortex are more extensively interconnected in KbDb KO mice. It is possible that not only the degree of connectivity differs, but also dendritic branching patterns and spine density, both of which are known to be regulated by neural activity and visual experience (Sin et al. 2002; Hofer et al. 2009). Modified rabies infection extensively and brightly labels neuronal processes, making it a useful method for analysis of morphology. Dendritic branching and spine density were therefore analyzed on apical and basal dendrites proximal to the soma of isolated GFP-labeled L2/3 pyramids (Fig. 2A, B). Dendrites were analyzed if a continuous length originating from the soma of ≥75 μm could be traced (average segment length: basal = 206 ± 13 μm; apical including side branches = 238 ± 22 μm). Dendritic branching complexity was assessed by reconstructing individual L2/3 pyramidal cells for Sholl analysis (Fig. 2A). The basal dendrites of KbDb KO-Tg cells had more complex branching (Fig. 2C; P < 0.0001; N = 9 WT-Tg and 11 KO-Tg cells), but the apical dendrites were indistinguishable from WT-Tg (Fig. 2D; P = 0.21; N = 9 WT-Tg and 10 KO-Tg cells).

Next, the postsynaptic components of excitatory synapses—the dendritic spines (Yuste and Bonhoeffer 2001; Nimchinsky et al. 2002)—were also analyzed along apical and basal dendrites of GFP-labeled L2/3 pyramids (Fig. 2B). Spine density along basal dendrites was increased in KbDb KO-Tg mice by about 10% over WT-Tg (Fig. 2E; WT-Tg: 6.8 ± 0.5 spines/10 μm, N = 11 cells, 1990 total spines; KO-Tg: density = 7.4 ± 0.3 spines/10 μm, N = 18 cells, 3784 total spines; P = 0.03). No difference in spine density was observed on apical dendrites (Fig. 2F; WT-Tg: 7.2 ± 0.8 spines/10 μm, N = 9 cells, 1365 total spines; KO-Tg: density = 7.7 ± 0.4 spines/10 μm; N = 12 cells, 2399 total spines; P = 0.84). There was no observed difference in spine type distribution along either basal or apical dendrites (Supplementary Fig. 3).

Bouton Density Is Elevated Along Axons Projecting to L2/3, and the Frequency of mEPSCs Recorded From L2/3 Pyramidal Cells Is Elevated, in KbDb KO Cortex

The increases observed in basal dendritic complexity and spine density in KbDb KO L2/3 pyramids imply that there might also be changes in presynaptic axons, such as an increase in the density of presynaptic axonal boutons. Small quantities of the anterograde tracer BDA were injected in the visual cortex to visualize axons of labeled cells (Fig. 3A). Bouton density along labeled axons running in L2/3 was significantly elevated in KbDb KO visual cortex compared with WT (WT: 6.7 ± 0.7 boutons/10 μm; KO: 11.0 ± 1.5 boutons/10 μm, P = 0.01; Fig. 3B,C). These experiments were performed in WT and KbDb KO mice. This approach not only provides information about axon bouton density, but also provides a measure of changes in connectivity independent of the transgenic line used above for the rabies tracing method.

To determine whether the increased density of both boutons and spines present in KbDb KO1 reflects a greater number of functional inputs, whole-cell recordings from L2/3 pyramids in WT and KbDb KO visual cortex were used to measure mEPSCs (Fig. 3D). KO mEPSC frequency was elevated relative to WT (WT: 1.4 ± 0.2 Hz; KO: 2.9 ± 0.6 Hz, P = 0.02; Fig. 3E), whereas mEPSC amplitude was similar between genotypes (WT: 19.7 ± 0.7 pA; KO: 18.9 ± 0.7 pA, P = 0.19; Fig. 3F). The presence of elevated mEPSC frequency is consistent with the idea that KO L2/3 pyramids receive more functional excitatory synaptic inputs than WT, whereas the similar mEPSC amplitudes suggests that the strength of individual synapses might not differ between genotypes.

Ocular Dominance Plasticity Is Increased in Both Juvenile and Adult Visual Cortex of KbDb KO Mice

Because there is a significant increase in cortical connectivity in KbDb KO mice present well past the critical period (P19–P32;
Gordon and Stryker 1996; Tagawa et al. 2005; Levelt and Hübener 2012), we wondered if there is also a change in OD plasticity that might even persist into adulthood. OD plasticity involves selective strengthening of synaptic inputs driven by the open eye and weakening of those driven by the closed eye in response to visual deprivation (Gordon and Stryker 1996), so an association between H2-Kb and H2-Db protein with synapses would be consistent with a role for these molecules in regulating plasticity. Therefore, we examined if H2-Kb and H2-Db protein is expressed in adult cortex. Previous work using antibodies of rather broad specificity showed that MHC1 protein is localized to neurons and at synapses (Corriveau et al. 1998; Huh et al. 2000; Datwani et al. 2009; Needleman et al. 2010). Here, newly generated and specific antibodies to H2-Kb and H2-Db were used to probe protein expression levels in western blots of synaptosome preparations in juvenile and adult cortex. Results using an H2-Kb antibody and an H2-Kb/Db antibody show that protein for these particular MHC1s can be detected in synaptosomes, implying that they are expressed at or near synapses both during the critical period (P28) and in adult (P100; Fig. 4A).

OD plasticity in KbDb KO mice was assessed at P100, months after the end of the visual cortical critical period in mice (Gordon and Stryker 1996; Tagawa et al. 2005). Multiple methods are available to assess OD plasticity, including induction of the immediate early gene Arc (Hofer et al. 2006a, 2006b). Arc is rapidly transcribed in neurons firing action potentials (Lyford et al. 1995), and Arc mRNA visualization can be used as a spatial readout of neurons functionally responding to open eye stimulation (Tagawa et al. 2005). To assess OD plasticity, at P100, mice received 10 days of monocular enucleation (ME) and subsequently at P110 Arc mRNA induction was used to assess OD plasticity at the L3/4 border in visual cortex. The width of the Arc mRNA in situ hybridization signal along the L3/4 border in visual cortex ipsilateral to the open (non-deprived) eye was measured in WT versus KbDb KO mice; this width is a reliable measure of open eye strengthening, a key component of OD plasticity (Frenkel and Bear 2004; Tagawa et al. 2005). Following 10 days of ME, the width of Arc mRNA in situ hybridization signal in KbDb KO mice is significantly greater than in WT (WT ME: 4 mice, 1147 ± 45 μm; KbDb KO ME: 6 mice, 1334 ± 32 μm; P = 0.02; Fig. 4B,C). Note that, in WT mice, there is
almost no detectable change in the width of Arc mRNA signal following 10 days of ME [WT normally reared (NR) vs. WT ME; \( P = 0.99 \)], consistent with the fact that the critical period ended months ago. Nor is there a difference between genotypes in NR mice not receiving visual deprivation (WT vs. KO NR; \( P = 0.9996 \); Fig. 4C). We also calculated a plasticity index, equal to the fold expansion of Arc signal following 10 days ME when compared with NR controls. The index is significantly greater for adult KbDb KO than for WT (WT: 1.01 ± 0.04, KbDb KO: 1.18 ± 0.03; \( P = 0.002 \); Fig. 4D), though it is lower than that seen during the critical period (from Datwani et al. 2009; WT: 1.51 ± 0.08, KbDb KO: 2.16 ± 0.11; \( P < 0.001 \)). Thus, in KbDb KO visual cortex, OD plasticity as assessed by measuring open eye strengthening appears to persist into adulthood.

**Discussion**

A major finding of this study is that pyramidal neuron local and horizontal connectivity in visual cortex of KbDb KO mice, as assessed both anatomically and functionally, is significantly greater than in WT. We examined connectivity after the end of the critical period and observed that excitatory connections in visual cortex of KO are increased over WT, particularly long distance horizontal connections hundreds of microns away. In addition, KbDb KO L2/3 pyramids have more complex basal dendrites as well as slightly elevated basal dendritic spine density and axon bouton density. mEPSC frequency of L2/3 pyramids is also elevated, suggesting that overall functional excitatory connectivity is greater in KO visual cortex. These observations demonstrate that one or both of these MHC class I proteins regulate the spatial pattern and the number of pyramidal cell connections in visual cortex. Furthermore, these data combined with previous findings (Datwani et al. 2009; Lee et al. 2014) suggest that H2-Db and/or Kb contribute specifically to the process of activity-dependent dendritic sculpting and synapse pruning that normally occurs during the visual cortical critical period.

**H2-Db and H2-Kb Regulate Intracortical Connectivity**

Neuronal MHC1 molecules were discovered in an unbiased screen for genes regulated by neural activity and visual experience (Corriuveau et al. 1998), and H2-Kb and H2-Db are known to be needed for developmental synapse remodeling (Datwani et al. 2009; Glynn et al. 2011; Elmer et al. 2013) and pruning of retinogeniculate projections (Lee et al. 2014). H2-Db and H2-Kb are expressed in cortical pyramidal neurons (Huh et al. 2000; Datwani et al. 2009; Needleman et al. 2010; Adelson et al. 2012), detected in cortical synaptosomes as shown here, and thought to be at the cortical synapse as viewed in electron micrographs (Needleman et al. 2010). Because H2-Kb and H2-Db are present in cortex, we suggest that the failure to remodel intracortical connections is due to loss of function in the cortex, rather than in the LGN or retina. This suggestion is underscored by the important observation that retinal activity, as well as visual acuity, in KbDb KO mice is indistinguishable from

**Figure 3.** Bouton density and mEPSC frequency are elevated in KbDb KO L2/3 pyramidal cells. (A–C) Anterograde tracing of cortical axons following a BDA injection into visual cortex. (A) Representative image of labeled axons in a projection site ipsilateral to the injection. Scale bar = 100 μm. (B) Higher magnification view of WT and KO axons in L2/3 used for analysis. Arrowheads indicate boutons. Scale bar = 10 μm. (C) Histograms showing bouton density in KbDb KO versus WT. WT: 6.7 ± 0.7 boutons/100 μm; KO: 11.0 ± 1.5 boutons/100 μm, \( P = 0.01 \). N = 9 WT and 9 KO animals; 1028 WT and 1078 KO boutons. (D–F) Whole-cell recordings from L2/3 pyramidal neurons in WT versus KO visual cortex slices. (D) Representative traces of WT and KO mEPSC recordings. (E and F) Cumulative histograms of mEPSC (E) frequency and (F) amplitude. Frequency: WT: 1.4 ± 0.2 Hz; KO: 2.9 ± 0.6 Hz, \( P = 0.02 \). Amplitude: WT: 19.7 ± 0.7 pA; KO: 18.9 ± 0.7 pA, \( P = 0.19 \). N = 26 WT cells from 6 animals and 29 KO cells from 7 animals. Each circle = one cell. *\( P < 0.05 \). Error bars = SEM. Mann–Whitney U-test. Analysis carried out at P40.
normal (Datwani et al. 2009; Lee et al. 2014). Thus, loss of neural activity, which might be expected to result in abnormal pruning, cannot explain the connectivity increases we observe here. Our observations are more consistent with the idea that Kb, Db, or both act downstream of activity as part of a pruning mechanism.

Results of our study correspond to and extend previous observations on a critical role for MHCI proteins in synapse elimination: In vitro, the frequency of mEPSCs and the density of synapses on rat cortical neurons can be modulated by expression of H2-Kβ (a mouse MHCI gene) or by using beta 2 microglobulin (β2m) siRNA (Glynn et al. 2011). In vivo in single β2m knockout or double β2m/TAP1 knockout mice, the frequency of mEPSCs is elevated in juvenile cortex, and the density of synapses is increased in both juvenile and adult cortex and hippocampus, as assessed using transmission EM (Goddard et al. 2007; Glynn et al. 2011). These provide useful albeit indirect evidence for a role for H2-Kβ or H2-Db in vivo specifically, because rat neurons express related but not identical MHC class I proteins to mice, and because β2m is an accessory molecule needed for stable surface expression of the majority (50+) of MHCI proteins (Zijlstra et al. 1990). In this study, we have examined mice lacking just 2 specific mouse MHCI molecules in vivo and observed not only an increase in functional synapses as assessed by mEPSC recordings, but also by directly examining spatial patterns of connectivity we have revealed an increase in excitatory pyramidal cell connectivity using the modified rabies virus transneuronal tracing method.

We further observed that dendritc complexity and spine density are elevated along proximal basal dendrites of L2/3 pyramids of KbDb KO visual cortex, but not along proximal apical dendrites. This selective effect on the basal dendrites of L2/3 cortical neurons in KbDb KO visual cortex is consistent with the fact that it is the basal rather than apical dendrites that receive the bulk of the intralaminar input from other L2/3 excitatory pyramids (Gilbert and Wiesel 1989; Thomson and Bannister 2003; Feldmeyer et al. 2006), which is transmitted via distinct holoreceptor complexes (neuropilin-2/plexin A3 regulates L5 pyramid spine distribution and dendritic arborization via distinct holoreceptor complexes (neuropilin-2/plexin A3 affects spine number vs. neuropilin-1/plexin A4 affects basal dendritic arborization; Tran et al. 2009). We suggest that H2-Db and H2-Kβ function to regulate basal dendritic complexity and spine density in L2/3 pyramidal neurons. However, we note that apical dendritic tufts in Layer 1 were not examined, and it is possible that H2-Db and H2-Kβ could also contribute to the complexity and spine density there, without affecting the proximal apical dendrite.

Greater Connectivity in KbDb KO Mice Could Serve as a Substrate for Ocular Dominance Plasticity and Recovery From Injury

Increased excitatory connectivity could underlie the observed enhancement of open eye strengthening in KbDb KO mice,
following eye closure both in the critical period and in adult. Previous work has shown that, in WT mice, eye closure generates new dendritic spines on neurons in primary visual cortex (Keck et al. 2008; Hofer et al. 2009), and retinal lesions that silence part of visual cortex result in the formation of new intralaminar axonal arbors (Yamahachi et al. 2009; Marik et al. 2013). In particular, new spines formed on L5 apical dendrites during MD persist even if binocular vision is restored; then a second MD later in life leads to more rapid and pronounced OD plasticity than normally observed in adult (Hofer et al. 2006a, 2006b), without recruiting any additional spines (Hofer et al. 2009). These results suggest that experience creates an anatomical substrate for functional plasticity that can be accessed later on.

In support of this hypothesis, mice lacking the MHCI receptor PirB (paired immunoglobulin-like receptor B) also have elevated spine density on the apical dendrites of L5 pyramidal neurons in visual cortex (Djurisic et al. 2013), and this increase is accompanied by enhanced OD plasticity not only during the critical period, but also in adulthood (Syken et al. 2006; Djurisic et al. 2013). Here, we demonstrate exuberant connectivity that persists in KbDb KO visual cortex past the close of the critical period, as well as enhanced OD plasticity in adulthood. Other molecules that restrict adult OD plasticity include Lynx1, chondroitin sulfate proteoglycans (CSPGs), and Nogo/NgR (Pizzorusso et al. 2002; McGee et al. 2005; Morishita et al. 2010; Miyata et al. 2012). If structural changes underlie functional plasticity, it would be interesting to determine if the absence of these molecules also is associated with anatomical correlates such as increased spine density or altered spatial patterns of connectivity.

Structural and functional plasticity are needed for recovery from brain injury. H2-Kb and H2-Db therefore could be useful therapeutic targets. For example, we have reported that KO of both H2-Kb and H2-Db is neuroprotective in an adult stroke model, where KO mice have less cell death and better behavioral recovery (Adelson et al. 2012) and knock out of PirB receptor protects Alzheimer’s model mice from memory loss (Kim et al. 2013). Defects in synapse pruning and plasticity during developmental critical periods have been hypothesized in disorders such as autism and Schizophrenia (Qu et al. 2011; Glausier and Lewis 2013), and it is noteworthy that the human MHC class I locus has been strongly correlated with schizophrenia in genome-wide association studies (Ripke et al. 2013, 2014; Stefansson et al. 2014). Our observations here in mouse visual cortex make a firmer link between developmental circuit refinement, plasticity, and neuronal MHCI class I function.

Authors’ Contributions

J.D.A. and C.J.S. designed experiments. J.D.A. conducted circuit tracing, spine analysis, and dendritic branching experiments with input and reagents from K.M. and L.L. J.D.A. and R.W.S. conducted axon tracing experiments. J.D.A., S.C., and M.D. performed OD plasticity experiments. H.L. recorded mEPSCs. B.K.B. performed biochemistry. J.D.A. and C.J.S. discussed and analyzed data and wrote paper.

Supplementary Material

Supplementary material can be found at: http://www.cercor.oxfordjournals.org/.

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References


